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- (57) Abstract

Screening procedures are disclosed for identifying compounds useful for inhibiting infection or pathogenicity. Methods are also disclosed for identifying pathogenic virulence factors.

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### METHODS OF SCREENING COMPOUNDS USEFUL FOR PREVENTION OF INFECTION OR PATHOGENICITY

#### Background of the Invention

The invention relates to screening procedures which identify compounds for inhibiting infection or disease in a eukaryotic host organism, or which induce or stimulate a host's pathogenic defense mechanisms. The invention also relates to the use of such compounds as anti-pathogens. In addition, the invention relates to procedures which identify pathogenic virulence factors.

Microbial pathogens such as bacteria, protozoa, fungi, nematodes, and viruses include a large and diverse group of organisms capable of infecting animals and plants. Initiation of an infection occurs when the infecting organism is pathogenic, and the host is susceptible to pathogenic invasion. After establishing contact with susceptible cells or tissues of the host, the pathogen acquires nutrients from its host,

- 20 facilitating its own survival. During the infection process the pathogen activates a cascade of molecular, biochemical, and physiological processes, the result of which is the release of substances detrimental to the host and the development of disease (See, e.g.,
- 25 Scientific American Medicine, W.H. Freeman and Co., CA, San Francisco, 1995; Agrios, G.N., Plant Pathology, Academic Press, 1988). The pathogenic effects of microbes are produced in a variety of ways.

Some pathogens act through secreted products.

30 Diphtheria, for instance, is caused by the bacillus,

Cornynebacterium diptheriae. This organism is inhaled by
the host and establishes infection in the upper
respiratory tract. While the bacterium does not itself
invade the bloodstream, its powerful toxins do. These

35 toxins are then absorbed by the cells of the body, enzyme
function is impaired, and host cells are destroyed.

25

Other diseases are the result of the body's reaction to a pathogen. For example, in pneumonia, a disease caused by Streptococcus pneumoniae, infection causes an outpouring of fluid and cells into the air sacs 5 of the lungs, interfering with respiration. Fungal infections of the skin similarly result from such inflammatory responses.

Yet other bacteria are opportunistic pathogens. Pseudomonas aeruginosa, for example, infects patients 10 with thermal burns and patients who are immunodeficient or otherwise immunologically compromised. P. aeruginosa infections can be acute and localized as in corneal ulcers and otitis media, chronic as in the lungs of cystic fibrosis patients, or systemic following 15 bloodstream invasion.

Plant pathogenic diseases are also of concern because they cause damage to plants and plant products. Phytopathogens produce disease in plants by any number of methods including: (1) consuming host cell nutrients; (2) 20 killing or disrupting host cell metabolism through toxins, enzymes, or growth-regulators; (3) affecting photosynthesis by inducing chlorosis (e.g., by degrading chloroplasts); and (4) blocking conductive tissues and interfering with normal physiological processes.

Crop plants, ornamentals, trees, and shrubs are especially vulnerable to diseases caused by bacteria, fungi, viruses, and nematodes. Phytopathogenic bacteria, for example, cause the development of many disease symptoms including leaf spots and blights, soft-rots, 30 wilts, overgrowths, scabs, and cankers. Bacterial diseases occur most commonly on vegetables (and some ornamentals) that have fleshy storage tissues, such as potatoes, carrots, onions, iris, or hyacinth. They may also occur in plants bearing fleshy fruit (such as 35 cucumber, squash, eggplant, or tomato), as well as in

leafy plants (such as cabbage, celery, lettuce, or spinach). Plant bacterial diseases occur throughout the world and cause serious damage to crops in the field, in transit, and in storage.

The mechanisms of plant pathogenesis are many and varied. One bacterial phytopathogen Erwinia, for example, causes plant diseases such as soft-rot and fire-blight by penetrating a plant through a wound or an accessible natural opening. Once inside, the bacteria secrete enzymes which break down the plant's middle lamellae, resulting in the maceration of tissue and ultimately cell death. Other bacteria, such as certain strains of Pseudomonas, may interfere with water translocation by disrupting xylem within the plant.

15 Pseudomonads invade the xylem of roots and stems and,

once inside, secrete enzymes and toxins which destroy the plant. Still other phytopathogenic bacteria, like Agrobacterium and Corynebacterium, stimulate cell division and cell enlargement in affected tissues. This generally leads to the development of amorphous overgrowths, galls, or tumors on roots, stems, or other organs (e.g., crown gall caused by Agrobacterium tumefaciens), or in the proliferation of infected organs (e.g., hairy root caused by Agrobacterium rhizogenes).

Prompt identification of the causative organism is essential to the appropriate selection of anti-pathogenic agents and successful management of clinical and agricultural infections. However, the extensive use of anti-pathogenic agents, such as sulfonamides,

30 tetracyclines, ampicillins, cephalosporins, and aminoglycosides, in both medicine and agriculture has strongly favored the selection of resistant microbial species. This is especially true of bacterial strains containing transmissible resistance plasmids. For

35 example, outbreaks of nosocomial infections from highly

resistant strains of Serratia, Klebsiella, Pseudomonas, Acinetobacter, Enterobacter, and Streptococcus have become important and recurrent problems. As a result of selecting resistant strains, over the past few decades, 5 P. aeruginosa has emerged as an important and problematic clinical pathogen, causing between 10% and 20% of infections in hospitals. Currently, several aminoglycosides and third-generation cephalosporins are efficacious against P. aeruginosa, but the relative ease with which P. aeruginosa acquires resistance necessitates the search for new compounds as potential replacements or alternative therapies.

#### Summary of the Invention

We have discovered that common pathogenic

virulence factors are involved in the infection and
pathogenicity of both animal and plant hosts. The
identification of such host-independent virulence factors
has facilitated improved screening methods designed to
evaluate and identify therapeutic agents useful for
inhibiting pathogenesis in either animal or plant hosts,
or both. Furthermore, our discovery provides the basis
for screening methods useful for identifying a variety of
new virulence factors. Identification of such virulence
factors also facilitates the development of targeted
reagents for use as anti-pathogens.

In a first aspect, therefore, the invention generally features a method for identifying a compound which is capable of inhibiting a pathogen in a eukaryotic host organism. The method involves (a) exposing (either sequentially or simultaneously) at least two different eukaryotic host organisms, at least one of the organisms being a non-rodent, to a single pathogen in the presence of at least one candidate compound; and (b) identifying a

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compound that inhibits the pathogen in each of the eukaryotic host organisms.

In preferred embodiments, the pathogen is a bacterium (e.g., Pseudomonas aeruginosa UCBPP-PA14); 5 the eukaryotic host organisms include a vertebrate (e.g., a non-rodent) and a plant, a vertebrate and an invertebrate; or an invertebrate and a plant. Preferably, the invertebrate is a nematode (e.g., a member of the genus Caenorhabditis); and the plant is a 10 crucifer (e.g., a member of the genus Arabidopsis). In other preferred embodiments, each of the eukaryotic host organisms is a plant; is a vertebrate; or is an invertebrate.

In a second aspect, the invention generally 15 features a method for identifying a compound which is capable of inhibiting a pathogen in a non-rodent eukaryotic host organism. The method involves (a) exposing a non-rodent eukaryotic host organism to a single pathogen in the presence of at least one candidate 20 compound; and (b) identifying a compound that inhibits the pathogen in the eukaryotic host organisms.

In one preferred embodiment, the pathogen is a bacterium (e.g., Pseudomonas aeruginosa UCBPP-PA14), and the non-rodent eukaryotic host organism is a nematode 25 (e.g., a member of the genus Caenorhabditis), and the plant is a crucifer (e.g., is a member of the genus Arabidopsis). In a second preferred embodiment, the pathogen is a bacterium (e.g., Pseudomonas aeruginosa UCBPP-PA14), and the non-rodent eukaryotic host organism 30 is a plant (e.g., is a member of the genus Arabidopsis).

In a third aspect, the invention generally features a method for identifying a pathogenic virulence factor. The method involves (a) identifying a pathogen which is capable of infecting at least two different 35 eukaryotic host organisms, at least one of the organisms

being a non-rodent; (b) generating a mutant of the pathogen; (c) exposing (either sequentially or simultaneously) each of the organisms to the mutated pathogen; (d) determining whether the mutated pathogen is capable of causing disease in each of the organisms, a reduction of disease in both of the organisms relative to that caused by the wild-type pathogen indicating a mutation in a pathogenic virulence factor; and (e) using the mutation as a marker for identifying the pathogenic virulence factor.

In a fourth aspect, the invention generally features a method for mutating a pathogenic virulence factor. The method involves: (a) identifying a pathogen which is capable of infecting at least two different eukaryotic host organisms, at least one of the organisms being a non-rodent; (b) generating a mutant of the pathogen; (c) exposing (either sequentially or simultaneously) each of the organisms to the mutated pathogen; and (d) determining whether the mutated pathogen is capable of causing disease in each of the organisms, a reduction of disease in both of the organisms relative to that caused by the wild-type pathogen indicating a mutation in a pathogenic virulence factor.

In a fifth aspect, the invention generally features a method of reducing the virulence of a pathogen. The method involves (a) identifying a pathogen which is capable of infecting at least two different eukaryotic host organisms, at least one of the organisms being a non-rodent; (b) generating a mutant of the pathogen; (c) exposing (either sequentially or simultaneously) each of the organisms to the mutated pathogen; and (d) determining whether the mutated pathogen is capable of causing disease in each of the organisms, a reduction of disease in both of the

organisms relative to that caused by the wild-type pathogen indicating a reduction in pathogen virulence.

By "inhibiting a pathogen" is meant the ability of a candidate compound to decrease, suppress, attenuate, 5 diminish, or arrest the development or progression of a pathogen-mediated disease or an infection in a eukaryotic host organism. Preferably, such inhibition decreases pathogenicity by at least 5%, more preferably by at least 25%, and most preferably by at least 50%, as compared to 10 symptoms in the absence of candidate compound in any appropriate pathogenicity assay (for example, those assays described herein). In one particular example, inhibition may be measured by monitoring pathogenic symptoms in a host organism exposed to a test compound or 15 extract, a decrease in the level of symptoms relative to the level of pathogenic symptoms in a host organism not exposed to the compound indicating compound-mediated inhibition of the pathogen.

By "non-rodent" is meant any organism that is not 20 a mouse, a rat, a guinea pig, or a hamster.

By "pathogenic virulence factor" is meant a cellular component (e.g., a protein such as a transcription factor) without which the pathogen is incapable of causing disease or infection in a eukaryotic host organism.

The invention provides long awaited advantages over a wide variety of standard screening methods used for distinguishing and evaluating the efficacy of a compound against microbial pathogens. For example, the screening methods described herein allow for the simultaneous evaluation of host toxicity as well as antipathogen potency in a simple in vivo screen. Moreover, the methods of the invention allow one to evaluate the ability of a compound to inhibit microbial pathogenesis, and, at the same time, to evaluate the ability of the

compound to stimulate and strengthen a host's response to pathogenic attack.

Accordingly, the methods of the invention provide a facile means to identify compounds that are safe for 5 use in eukaryotic host organisms (i.e., compounds which do not adversely affect the normal development and physiology of the organism), and efficacious against pathogenic microbes (i.e., by suppressing the virulence of a pathogen). In addition, the methods of the 10 invention provide a route for analyzing virtually any number of compounds for anti-pathogenic effect with highvolume throughput, high sensitivity, and low complexity. The methods are also relatively inexpensive to perform and enable the analysis of small quantities of active 15 substances found in either purified or crude extract Furthermore, the methods disclosed herein provide a means for identifying anti-pathogenic compounds which have the capability of crossing eukaryotic cell membranes and which maintain therapeutic efficacy in an in vivo 20 method of administration.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

#### Detailed Description

The drawings will first be described.

#### Drawings

25

rigure 1 is a photograph showing the symptoms caused by Pseudomonas syringae and Pseudomonas aeruginosa on Arabidopsis (ecotype Llagostera (Ll)) leaves. Mockinoculated (left); Pseudomonas syringae pv. maculicola strain ES4326 (center); Pseudomonas aeruginosa strains UCBPP-PA14 (right).

Figures 2A-D are graphs showing the growth of Pseudomonas syringae and Pseudomonas aeruginosa in

Arabidopsis leaves. Figure 2A is a graph showing the growth of Pseudomonas syringae pv. maculicola strain ES4326 (open squares), Pseudomonas aeruginosa strain UCBPP-PA14 (open circles), and Pseudomonas aeruginosa 5 strain UCBPP-PA29 (open triangles) in ecotype Llagostera. Figure 2B is a graph showing the growth of Pseudomonas aeruginosa strain UCBPP-PA14 in three Arabidopsis ecotypes: Columbia (solid squares); Argentat (solid circles); and Bensheim (solid triangles). Figure 2C is a 10 graph showing the growth of Pseudomonas aeruginosa strain UCBPP-PA14 (solid circles) and isogenic plcS (open squares), and toxA (open diamonds) mutants. Figure 2D is a graph showing the growth of Pseudomonas aeruginosa strain UCBPP-PA14 (solid circles), isogenic gacA (open 15 diamonds), and degP (open squares) mutants in ecotype Llagostera. Bacterial counts in Arabidopsis leaves were performed as described herein. Means of four samples ± SD are shown. Three independent experiments gave similar results. Incubation conditions for the plants were 20 identical to the experiments presented in Table I, infra.

Figure 3 is a graph showing a comparison of Caenorhabditis elegans lethality growing on wild-type Pseudomonas aeruginosa strain UCBPP-PA14 and on an isogenic degP mutant.

Pigure 4 is a graph showing a comparison of Caenorhabditis elegans lethality growing on wild-type Pseudomonas aeruginosa strain UCBPP-PA14 and on an isogenic gacA mutant.

Below we describe experimental evidence

30 demonstrating that a bacterial pathogen is capable of
causing disease in both a plant, in an animal, and in a
nematode, and that there is an overlap in virulence
factors responsible for causing microbial pathogenic
disease in plants, animals, and nematodes. These

experimental examples are intended to illustrate, not limit, the scope of the claimed invention.

Identification of Common Virulence Factors Required for Pseudomonas aeruginosa Pathogenicity in Plants and

#### 5 Animals

To identify multi-host virulence factors, we first searched for bacterial pathogens capable of eliciting disease in both plant and animal pathogenesis models. A variety of P. aeruginosa isolates were screened using an Arabidopsis thaliana leaf pathogenesis infiltration system. Isolates which elicited disease symptoms in Arabidopsis were then tested for pathogenicity in a mouse full-thickness skin burn model and a nematode feeding assay.

Specifically, we first screened a collection of 15 P. aeruginosa strains which included 30 human clinical isolates, 20 soil isolates, and 25 plant isolates (obtained from the University of California at Berkeley, Department of Plant Pathology). Each of these isolates 20 was independently injected into the leaves of four different Arabidopsis ecotypes (land races or wild accessions) to determine whether the isolate was a plant pathogen. Several Arabidopsis ecotypes were assayed to increase the likelihood of identifying a suitable 25 pathogen because plant pathogens, including Arabidopsis pathogens, typically exhibit a high level of host cultivar or ecotype specificity. Multiple host assays were also carried out because P. aeruginosa strains exhibiting ecotype specificity were more likely to be 30 bona fide plant pathogens (rather than artifactual pathogens, capable of infecting plants only in the artificial environment created in the laboratory).

Screening experiments using an Arabidopsis leaf pathogenesis infiltration system were performed as

follows. P. aeruginosa strains were grown in Luria Broth (LB) medium at 37°C, washed twice in 10 mM MgSO<sub>4</sub>, resuspended at an optical density of 600 [OD 600] = 0.2 in 10 mM MgSO<sub>4</sub>, diluted 1:100 (corresponding to a bacterial 5 density of 103 cfu/cm2), and injected into leaves of six week old Arabidopsis plants. Plants were kept in a growth chamber during the course of the experiment at 28-30°C and 90-100% relative humidity. Disease symptoms and growth were monitored daily for five days. 10 elicited five days post-injection were characterized as: "none," no symptoms; "weak," localized weak water-soaking and chlorosis (yellowing) of tissue circumscribing the injection site; "moderate," moderate water-soaking and chlorosis with the majority of tissue softened around the 15 inoculation site; or "severe," severe soft-rotting of the entire inoculated leaf characterized by a water-soaked reaction zone and chlorosis circumscribing the injection site at 2-3 days post-injection. The soft-rot symptoms pervaded the leaf at 4-5 days post-injection. 20 intercellular fluid containing bacteria was harvested at five days, and bacterial counts were determined according to standard methods (see, e.g., Dong et al. (1991) Plant Cell 3:61). Four different samples were taken using two leaf discs per sample. Three independent experiments 25 gave similar results. Control plants inoculated with 10 mM MgSO<sub>4</sub> showed no symptoms during the course of the experiments. In other control experiments, none of the genetically characterized P. aeruginosa strains PAK, PAO1, or PO37 caused appreciable symptoms on any of the 30 Arabidopsis ecotypes tested. These strains were found to be non-pathogenic in the ecotypes tested, but pathogenic in culture.

While the majority of the 75 P. aeruginosa strains which were screened caused no symptoms in Arabidopsis

35 leaves, several strains elicited weak to moderate soft-

rot symptoms characterized by chlorosis and water-soaking of the tissue circumscribing the injection site. strains, UCBPP-PA14 (a human clinical isolate) and UCBPP-PA29 (a plant isolate) caused severe soft-rot symptoms in 5 some of the ecotypes tested, typical of a highly virulent Table I shows the growth of plant bacterial pathogen. P. aeruginosa UCBPP-PA14 and UCBPP-PA29 five days post infection, and disease symptoms elicited by these P. aeruginosa strains on different Arabidopsis ecotypes. 10 particular, strain UCBPP-PA14 caused severe soft-rotting in both the Llagostera (L1) and Columbia (CO1) Arabidopsis ecotypes, but caused no symptoms in ecotype Argentat (Ag) and only moderate symptoms in ecotype Bensheim (Be). Table I also illustrates that strain 15 UCBPP-PA29 caused severe symptoms in Ll and weak symptoms in Col, but caused no symptoms in Ag or Be.

P.aeruginosa UCBPP-PA14		TABLE I P.aeruginosa UCBPP-PA29	inosa PA29	
Arabidopsis Ecotype	Arabidopsis cfu/cm²leaf area Ecotype	Symptoms	cfu/cm2leaf area	Sympt
Llagostera	Llagostera 2.6 x 107±2.0 x 107 Severe	Severe	2.7 x 10 <sup>7</sup> ±1.3 x 10 <sup>7</sup> Sever	Sever
Columbia	9.0 x 10 <sup>6</sup> ±6.0 x 10 <sup>6</sup> Severe	Severe	6.0 x 105±3.0 x 105 Weak	Weak
Argentat	3.0 x 10 <sup>5</sup> ±1.4 x 10 <sup>5</sup> None	None	1.5 x 10 <sup>5</sup> ±9.0 x 10 <sup>4</sup> None	None
Bensheim	1.1 x 10 <sup>6</sup> ±4.9 x 10 <sup>5</sup> Hoderate		4.5 x 105±2.0 x 105 None	None

As shown in Figure 1, the severe symptoms elicited by UCBPP-PA14 (far right) were characterized by a water-soaked reaction zone and chlorosis, resulting in complete maceration and collapse of the leaf 4 to 5 days post-infection (compare with control far left). These symptoms were essentially indistinguishable from the symptoms elicited by the highly virulent Arabidopsis pathogen Pseudomonas syringae pv. maculicola strain ES4326 (pictured in center).

To verify that the severity of disease symptoms 10 was correlated with bacterial proliferation, growth of each of the strains UCBPP-PA14 and UCBPP-PA29 was measured over the course of several days in Arabidopsis leaves as described above. As shown in Figure 2A, 15 strains UCBPP-PA14 (open circles) and UCBPP-PA29 (open triangles) reached maximal bacterial density of approximately 107 cells/cm2 leaf area by five days in ecotype Ll, which corresponded to 104-fold increases from the initial inocula. The growth profiles of these 20 strains in Ll was similar to that of the virulent Arabidopsis pathogen P. syringae pv. maculicola strain ES4326 (Figure 2A, open squares). Strain UCBPP-PA14 also proliferated 104-fold in ecotype Col (Figure 2B, solid squares; Table I). In contrast, strain UCBPP-PA14 25 increased only 103- and 102-fold in Be and Ag leaves, respectively (Figure 2B, solid triangles and solid circles, respectively; Table I), and strain UCBPP-PA29 increased only  $10^2$ - to 6 x  $10^2$ -fold in ecotypes Col, Ag, and Be (Table I). In each case, reduced bacterial counts 30 in leaves reflected less severe symptom development. Accordingly, each of these P. aeruginosa strains was similar to other phytopathogenic bacteria in its ability to cause disease in an ecotype-specific manner.

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UCBPP-PA14 and UCBPP-PA29 isolates found to elicit disease symptoms in Arabidopsis were then tested in a mouse full-thickness skin burn injury assay. This involves 5% of the murine body surface area fashioned on an outstretched area of abdominal skin (Stevens et al. (1994) J. of Burn Care and Rehabil. 15:232). In this model, the damaged epidermis and dermis undergoes coagulation necrosis, but the underlying rectus abdomini (RA) muscles are not injured. In the absence of infection, all animals survive.

aeruginosa inoculum is injected intradermally into the midline crease of the burn eschar. The bacteria proliferate in the burn wound, and some strains may invade the normal underlying RA muscles. Highly pathogenic strains can also invade the vasculature. The number of bacteria found in the RA muscles underlying and adjacent to the burn after 24 hours gives a quantitative measure of local invasiveness, and mortality indicates both local and systemic invasiveness.

Mouse full-thickness skin burn studies were performed as follows. Six week old male CD-1 mice (Charles River Animal Farms) weighing between 25 and 35 grams were used in all experiments, following an animal 25 burn model (Stevens et al., supra). Mice were injected with ~5 x 10<sup>3</sup> cells. No viable bacterial cells were retrieved from the underlying RA muscle immediately after bacterial injection or in animals who received a sham injury in other studies. In mortality studies,

30 immediately following the burn, mice were injected with 10<sup>2</sup> cells, and the number of animals which died of sepsis was monitored each day for ten days. Two groups of control animals consisting of (i) mice burned but not injected and (ii) mice injected with heat-killed UCBPP-35 PA14 resulted in 0% mortality.

Data shown in Table II (below) illustrate the proliferation of P. aeruginosa strains in a mouse full-thickness skin burn model. Table II indicates that strains UCBPP-PA14 and UCBPP-PA29 proliferated and invaded the RA muscles comparably to the well-characterized P. aeruginosa human isolates PO37, PAK, and PA01. All strains reached titers ranging from 1.8 x 108 to 3.6 x 108 cfu per gram tissue in RA muscle biopsies taken directly beneath the burn and infection site (Table II). Furthermore, all strains reached titers ranging from 4.0 x 107 to 8.2 x 107 cfu per gram tissue in RA muscle biopsies taken adjacent to the burn. In addition, tissue samples processed for routine histology revealed that strain UCBPP-PA14 invaded the muscle to the same degree as strain PO37.

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#### TABLE II

	P. aeruginosa Strain	Mean titer ± S.I in biopsies underneath burn	Mean titer ± S.D. in biopsies adjacent to burn
5	UCBPP-PA14	$20.0 \times 10^{7} \pm 9.0 \times 10^{7}$	$6.0 \times 10^7 \pm 2.1 \times 10^7$
	UCBPP-PA29	$36.0 \times 10^7 \pm 10.0 \times 10^7$	$8.2 \times 10^{7} \pm 2.0 \times 10^{7}$
	PO37	$30.0 \times 10^7 \pm 11.0 \times 10^7$	$5.8 \times 10^{7} \pm 1.0 \times 10^{7}$
	PAK	$18.0 \times 10^{7} \pm 9.1 \times 10^{7}$	$6.0 \times 10^7 \pm 1.2 \times 10^7$
	PAO1	$31.0 \times 10^7 \pm 10.0 \times 10^7$	$4.0 \times 10^{7} \pm 1.8 \times 10^{7}$

in comparison to PO37 was also assessed by conducting mortality studies in the mouse full-thickness skin burn model as described above. Strains UCBPP-PA14, UCBPP-PA29, and PO37 caused 77% (17/22), 6% (1/16), and 22% (2/9) mortality, respectively, by the tenth day post-burn and infection (Table III). Additional experiments showed strains PA01 and PAK caused significantly less mortality in this model than UCBPP-PA14.

Strain UCBPP-PA14 was then selected for additional 20 studies because it was infectious in both plant and animal pathogenicity models in which the outcome of pathogenesis could be quantitated, and because the level of virulence in these models was comparable to known plant and animal pathogens. Specifically, we sought to 25 determine whether there were common virulence determinants in strain UCBPP-PA14 required for pathogenicity in both hosts. Our strategy was to use a marker exchange procedure to generate UCBPP-PA14 mutants carrying insertion mutations in four different genes, two 30 known to be virulence determinants for P. aeruginosa in animal hosts, one known to be a virulence determinant for phytopathogenic bacteria in plant hosts, and one known to be a virulence determinant for several animal bacterial pathogens in animal hosts. The two animal virulence 35 genes of P. aeruginosa were plcs and toxA encoding the

exported proteins phospholipase C and exotoxin A, respectively (Ohman et al. (1980) Infect. Immun. 28: 899; Ostroff et al. (1987) J. Bacteriol. 169: 4597). Exotoxin A ribosylates G proteins, and phospholipase C 5 preferentially degrades phospholipid of eukaryotic cells (Iglewski et al. (1975) Proc. Natl. Acad. Sci. 72:2284; Berka et al. (1982) J. Bacteriol. 152:239). The plant pathogen virulence determinant was gacA, identified as a global regulator of excreted anti-fungal factors in the 10 non-pathogenic soil bacterium P. fluorescens (Laville et al. (1992) Proc. Natl. Acad. Sci. 89:1562; Gaffney et al. (1994) Mol. Plant-Microbe Interact. 7:455). In the phytopathogens P. syringae pv. syringae and P. cichorii, gacA appears to serve as a transcriptional regulator of 15 genes that encode extracellular products involved in pathogenicity (Rich et al. (1994) J. Bacteriol. 176:7468). The other animal virulence determinant, degP (also known as htrA), has been identified as a stressresponse protease which is responsible for degrading 20 incorrectly folded periplasmic proteins in Brucella and Salmonella (Elzer et al. (1994) Infection and Immunity 62: 4135; Johnson et al. (1991) Mol. Microbiol. 5: 410). The UCBPP-PA14 homologues of plcs and toxA were identified in a genomic cosmid library of strain UCBPP-25 PA14 using cloned DNA fragments corresponding to the plcs and toxA genes of P. aeruginosa strain PAK as hybridization probes. A genomic library of strain UCBPP-PA14 was prepared according to standard methods in the cosmid cloning vector pJSR1, which was itself constructed 30 by ligating a 1.6 kb BglII fragment containing the bacteriophage lambda cos site from pHC79 (see, e.g., Hohn et al. (1980) Gene 11: 291) into the BglII site of pRR54 (see, e.g., Roberts et al. (1990) J. Bacteriol. 172:

6204). A 1.7 kb BamH1 fragment isolated from plasmid 35 pMS150 containing the toxA gene (see, e.g., Lory et al.

(1983) Gene 22:95) and a 3.0 kb BamH1-PstI fragment isolated from plasmid pSL2 (see, e.g., Lory et al. (1988) J. Bacteriol. 170:714) containing the plcS gene were used to probe the UCBPP-PA14 genomic library in pJSR1.

The UCBPP-PA14 homologue of gacA was identified in the same cosmid library using a PCR-amplified product corresponding to a conserved region of the P. fluorescens gacA gene according to standard methods. The oligonucleotides 5'- GCTAGTAGTCGATGACC- 3' (SEQ ID NO:1)

10 and 5'-GCTGGCATCAACCATGC- 3' (SEQ ID NO:2) were designed on the basis of the sequence of the gacA gene (Laville et al. (1992) Proc. Natl. Acad. Sci. 89:1562) and used to amplify a 625 base-pair product containing the gacA gene of Pseudomonas fluorescens, which in turn was used to probe the UCBPP-PA14 genomic library in pJSR1 described above. The UCBPP-PA14 homologue of the degP gene was identified in the UCBPP-PA14 cosmid library using the degP gene of Pseudomonas syringae pv. maculicola as a probe.

All four genes were subcloned and mutagenized by the insertion of a cassette encoding gentamicin resistance using standard methods.

In addition, a 6 kb BamHI fragment isolated from the cosmid clone containing the plcS gene of strain

25 UCBPP-PA14 was subcloned from a pJSR1-derived cosmid into the BamHI site of pBR322. The resulting clone, pLGR101, was mutagenized by insertion of a gentamicin-encoding DNA cassette into the XhoI site of the plcS gene to construct pLGR201. The gentamicin-resistance gene cassette is a

30 1.8 kb BamHI fragment from plasmid pH1JI (see, e.g., Rubin (1987) Plasmid 18, 84). A 1.6 kb BamHI fragment containing the toxA gene was subcloned from a pJSR1-derived cosmid into pBR322 to construct pLGR102 and subsequently mutated by introducing the gentamicin

35 cassette into the BglII site of the toxA gene to

construct plasmid pLGR202. And a 2.5 kb HindIII-EcoRI fragment containing the P. aeruginosa strain UCBPP-PA14 gacA gene was subcloned from a pJSR1-derived cosmid into pBR322 to construct pLGR103. The presumptive gacA gene 5 was partially sequenced to confirm that the UCBPP-PA14 gacA had been cloned. pLGR103 was mutagenized by inserting the gentamicin cassette into the SalI site of gacA to construct the plasmid pLGR203. A 1.6 Pst I fragment containing part of the degP gene was subcloned 10 from pPY201 a derivative of the cosmid clone pH126 of the strain UCBPP-PA14 into the PstI site of pUC19 to construct pNAS. A 1.6 kb Sall fragment containing the gentamicin cassette was inserted into the XhoI site of the degP gene in pNAS to construct pNASGm. Next, a 3.2 15 kb SphI/XhoI fragment was isolated from the pNASGm vector and subcloned into the SphI/XhoI sites of pCVD442 to construct pPY206, which contained the mutated degP gene.

The mutated genes were transferred to the UCBPP-PA14 genome using standard marker exchange techniques, and the structures of the resulting marker exchange mutations were verified by DNA blot analysis. Thus, plasmids pLGR201, pLGR202, pLGR203, and pPY206 were used for gene replacement of the plcs, toxA, gacA, and degP genes respectively, by the method described in Rahme et al. (1991) J. Bacteriol. 170:575, using gentamicin at 30 mg/mL to screen for the double crossover events and carbenicillin at 300 mg/mL to screen for the loss of the vector. None of these four mutations had any detectable effect on the growth of the bacteria compared to wild-type in either rich or minimal media.

The effects of the plcs, toxA, gacA, and degP mutations on the pathogenicity of UCBPP-PA14 in the Arabidopsis model were tested by infiltrating the mutant strains into Arabidopsis ecotype Ll. Unlike wild-type UCBPP-PA14, none of the mutants caused maceration or

mutant caused attenuated soft-rot and chlorosis symptoms without the accompanying maceration of the affected tissue characteristic of UCBPP-PA14. The plcs, gacA, and degP mutants elicited even weaker symptoms, causing only chlorosis. Consistent with the attenuated symptoms, growth of the toxA, plcs, gacA, and degP mutants after 5 days was approximately 10-fold, 10<sup>2</sup>-fold, 5 x 10<sup>3</sup>-fold, and 10<sup>2</sup>-fold less, respectively, than the growth of the wild type (Figures 2C and 2D).

The growth and symptoms of the three mutants tested (plcS, toxA, and gacA) were fully restored to wild-type levels in plants when these mutants were complemented with the corresponding wild-type genes 15 carried on a plasmid. This was accomplished by subcloning a 6 kb BamH1 fragment from the cosmid clone pB85 of the genomic library containing the plcSR operon of strain UCBPP-PA14 into the BamHI site of plasmid pRR54 to construct pLGR301. Plasmid pLGR301 was then used for 20 the genetic complementation studies of the plcs mutant. A 2.4 kb EcoRI/EcoRV fragment isolated from plasmid pMS150 containing the toxA gene of the strain PAK, was subcloned into the EcoRI/EcoRV sites of plasmid pBR322 to construct pLGR106. From pLGR106 a SphI/PstI fragment 25 containing toxA was cloned into the SphI/PstI sites of pRR54 to construct pLRG206. A 1.2 kb HindIII/XhoI fragment containing the gacA gene was isolated from cosmid clone pH106 and subcloned into the HindIII/SalI sites of plasmid pRR54 to construct pLGR204. Plasmids 30 pLRG206 and pLGR204 were then used for genetic complementation studies of the toxA and gacA mutants.

Table III shows lethality studies corresponding to these three mutant P. aeruginosa strains in a mouse full-thickness skin burn model. In such lethality studies, 35 mice that were burned and infected with either plcs or

toxA mutants exhibited significantly lower mortality (40%
with both mutants) compared to infection with the wildtype strain (77%). The gacA and degP mutants caused no
mortality (Table III). The differences in mortality

5 rates between the mutants and wild-type was statistically
significant at the 95% or greater confidence level.
Statistical significance for mortality data was
determined by using the chi-square test with Yates'
correction. Groups were considered statistically

10 significant at P ≤ 0.05. All the mutants achieved
statistical significance (plcS and toxA, P = 0.05; gacA,
P = 0.00005).

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#### TABLE III

P. aeruginosa Strain	Mortality ratio of mice at 10 days following burn and infection
UCBPP-PA14	17/22
UCBPP-PA14 plcs	6/15
UCBPP-PA14 toxA	6/15
UCBPP-PA14 gack	0/10
UCBPP-PA14 degF	0/11
UCBPP-PA29	1/16
PO37	4/9

10

The above results demonstrate that plcs, toxA, gacA, and degP are involved in both plant and animal pathogenesis and indicate a part of the pathogen's machinery required for disease development is common or shared in animal and plant hosts. One of the shared virulence factors, gacA, is active at the regulatory level, demonstrating that mechanisms for regulation of virulence factors are conserved between plant and animal pathogens. The plcS and toxA gene products are specific virulence determinants which presumably attack the membranes and inhibit protein synthesis in both plant and animal cells, respectively.

the pathogenicity of P. aeruginosa UCBPP-PA14 was
25 measured in a nematode feeding assay. The feeding assay
was set up as follows. First, 5 µl of an overnight
culture of P. aeruginosa UCBPP-PA14, or an isogenic
strain of P. aeruginosa UCBPP-PA14 carrying a degP or
gacA mutation, was inoculated onto the center of an NGM
30 agar plate and cultured for 24 hours at 37°C. After
several hours of cooling at room temperature, the plates
were seeded with eight Caenorhabditis elegans L4-stage
worms. Plates were subsequently incubated in the dark at
25°C, and deceased worms were scored every 6 hours. A

worm is considered dead when it is non-motile, no longer displays any pharyngeal pumping action, and no longer exhibits defecation behavior.

Figures 3 and 4 show the results of the nematode 5 feeding lethality assay using wild-type UCBPP-PA14 and its degP and gacA isogenic mutants, respectively. results depicted in both Figure 3 and Figure 4 show that P. aeruginosa UCBPP-PA14 kills C. elegans. The results also show that isogenic mutants of P. aeruginosa UCBPP-10 PA14 carrying insertions which functionally disabled either the degP or gacA gene were significantly reduced in virulence in both the nematode and mouse fullthickness skin burn assay (Figures 3 and 4; Table III). The gacA gene is known to be a virulence determinant for 15 P. syringae in plant hosts, and degP is known to be a virulence factor for both P. syringae and Salmonella typhimurium. As is discussed below, we have used these screening methods for identifying several mutants that exhibit reduced pathogenicity in nematodes and 20 Arabidopsis; three of the mutants we isolated were found to be less pathogenic in mice.

described herein has several practical ramifications.

For example, these results indicate the molecular basis
of pathogenesis is remarkably similar in plants and animals. Thus, as described below, the multi-host pathogen system can be used for the identification and study of new virulence factors. In particular, the entire P. aeruginosa genome can be scanned for pathogenicity-related genes by testing individually mutagenized P. aeruginosa in different host organisms, e.g., using the Arabidopsis or nematode assays described herein. Genes identified in this manner can then be tested in the mouse full-thickness skin burn model. This system also facilitates the elucidation of the molecular

basis of host specificity of bacterial pathogens.

Virulence factors identified using this model system

provide targets for the development of a new generation

of chemical therapies for both clinical and agricultural

microbial diseases.

Screening Systems For Identifying Common Virulence Genes

Based on the results described above showing that a set of P. aeruginosa virulence factors are involved in pathogenicity in three diverse hosts and that these 10 common virulence determinants define fundamental features of bacterial pathogenicity which are host independent, we have developed a method for identifying virulence determinants important for pathogenicity in plants and animals. The screen utilizes a multi-host animal/plant 15 pathogen (e.g., P. aeruginosa UCBPP-PA14) and exploits the ability to readily screen thousands of randomly generated microbial mutants in virtually any host organism. Useful eukaryotic host organisms include, without limitation, nematodes (e.g., Caenorhabditis 20 elegans), plants (e.g., a seed or leaf from Arabidopsis), yeast or other fungi, fish (e.g., zebrafish), flies (e.g., Drosophila melanogaster), mice, and the like. general, a microbial pathogen is mutated according to standard methods known in the art and then subsequently 25 evaluated for its ability to induce disease in the host organism. Mutagenized pathogens found to have diminished pathogenicity or which are rendered non-pathogenic are useful in the method of the invention. Such mutant pathogens are then used for identifying host-dependent or 30 host-independent virulence factors responsible for pathogenicity according to methods known in the art.

The following is a working example of a virulence factor nematode screening system which utilizes the human clinical isolate P. aeruginosa UCBPP-PA14 found to be

infectious in three different models: a mouse skin fullthickness burn model, a C. elegans nematode feeding model, and an Arabidopsis thaliana leaf infiltration model. The advantage of using a nematode as a host for 5 studying a human or plant pathogen such as Pseudomonas is the relative simplicity of identifying non-pathogenic Pseudomonas mutants in the nematode. For example, a C. elegans screen consists of putting two L4 stage worms on a lawn of a P. aeruginosa mutant and looking for 10 surviving worms after 5 days. A pathogen such as P. aeruginosa UCBPP-PA14 is mutated according to any standard procedure, e.g., standard in vivo or in vitro insertional mutagenesis methods (see, e.g., Kleckner et al. (1977) J. Mol. Biol. 116:125). Other methods are 15 also available, e.g., chemical mutagenesis. By the fifth day, very few or no live worms can be found in the plate seeded with wild-type, pathogenic bacteria, whereas on a plate with E. coli or a non-pathogenic mutant, hundreds or thousands of live progeny of the initial two 20 hermaphrodite worms are present. Thus, worms growing in the presence of mutated P. aeruginosa is an indication that a gene responsible for pathogenicity has been The positions of an inactivating mutations inactivated. are mapped, leading to the cloning and identification of 25 the mutated virulence factor (e.g., by nucleotide sequencing).

To identify genes involved in pathogenicity, we generated mutants of P. aeruginosa UCBPP-PA14 using standard techniques of transposon mutagenesis (see, e.g. 30 Manoil et al. (1985) Proc. Natl. Acad. Sci. 82:8129; Taylor et al. (1989) J. Bacteriol. 171:1870); over 8000 mutants were generated. The pathogenicity of 1900 of these mutants was then assessed using the C. elegans feeding assay described above. As shown in Table IV, we

isolated 8 UCBPP-PA14 mutants that exhibited attenuated pathogenicity in C. elegans.

In addition, we also examined the pathogenicity of another collection of mutants generated by transposon

5 mutagenesis in a lettuce leaf pathogenesis assay using standard methods (see, e.g., Cho et al. (1975)

Phytopathology 65:425). Using this assay, we isolated 2900 UCBPP-PA14 mutants with attenuated pathogenicity on lettuce leaves. These mutants were subsequently tested in the Arabidopsis leaf pathogenesis assay according to the methods described herein. As shown in Table IV, we isolated 12 UCBPP-PA14 mutants that exhibited attenuated pathogenicity in Arabidopsis.

TABLE IV

		Arabidopsis thaliana	C. elegans
15	No. of mutants tested	2900	1900
4	No. of attenuated mutants	12	8

One UCBPP-PA14 mutant identified in the Arabidopsis infiltration assay was then tested for pathogenicity in both the C. elegans feeding assay and the mouse full-thickness skin burn assay. We found that this UCBPP-PA14 mutant was less pathogenic in both systems when compared to the wild-type UCBPP-PA14 strain. Furthermore, we also tested two mutants identified in the Arabidopsis bioassay for pathogenicity in the mouse full-thickness burn assay. These mutants were also found to be less pathogenic in mice when compared to the wild-type UCBPP-PA14 strain. Together these results provide further evidence for the existence of common virulence factors for pathogenicity in plants and animals.

The results described above demonstrate that pathogenic interactions occur between P. aeruginosa UCBPP-PA14 and C. elegans. Strain UCBPP-PA14 kills C.

elegans. UCBPP-PA14 is also infectious in an Arabidopsis thaliana leaf infiltration assay (Figures 1 and 2; Table I) and in a mouse full-thickness skin burn model (Tables II and III). Furthermore, we have demonstrated that null mutations in UCBPP-PA14 degP and gacA genes significantly decrease pathogenesis in all three models. Thus, we have provided the first evidence for the existence of common virulence factors for pathogenicity in plants and animals. Such virulence factors make possible the isolation of compounds that interfere with virulence factor function (e.g., through direct reduction of pathogenicity or enhancement of a host response), and also make possible the identification of these compounds in simple experimental systems (e.g., Caenorhabditis).

# 15 <u>Screening Systems for Identifying Therapeutics or Plant</u> Protectants

As discussed above, our experimental results demonstrate that a set of P. aeruginosa virulence factors are involved in pathogenicity in three diverse hosts and 20 that these common virulence determinants define fundamental features of bacterial pathogenicity which are host independent. Based on this discovery we have developed a screening procedure for identifying therapeutic compounds (e.g., anti-pathogenicity 25 pharmaceuticals) which can be used to inhibit pathogens capable of independently infecting either an animal (e.g., a human patient) or a plant (e.g., a commercial crop plant). In general, the method involves screening any number of compounds for therapeutically- or 30 agriculturally-active agents by employing the multi-host animal/plant pathogen (e.g., P. aeruginosa UCBPP-PA14) system(s) described herein. Based on our demonstration that there are common virulence factors for pathogenicity in plants, mice, and nematodes, it will be readily

understood that a compound which interferes with the function of such a virulence factor in a nematode also provides an effective therapeutic agent in a mammal (e.g., a human patient) or a plant. Whereas most antibiotics currently in use in medicine or agriculture are either bactericidal or bacteriostatic, thus favoring strains or mutants resistant to them, the compounds identified in the screening procedures described herein (e.g., the nematode system) do not kill the bacteria but instead render them non-pathogenic. Moreover, since the screening procedures of the invention are performed in vivo, it is also unlikely that the identified compounds will be highly toxic to a eukaryotic host organism.

Accordingly, the methods of the invention simplify
the evaluation, identification, and development of active
agents such as drugs and plant protectants for the
treatment of pathogenic diseases, including diseases
caused by bacteria, fungi, viruses, annelids, nematodes,
platyhelminthes, and protozoans. In general, the
screening methods of the invention provide a facile means
for selecting natural product extracts or compounds of
interest from a large population which are further
evaluated and condensed to a few active and selective
materials. Constituents of this pool are then purified
and evaluated in the methods of the invention to
determine their anti-pathogenic activity.

Below we describe screening methods for evaluating the efficacy of a compound as an anti-pathogenic agent.

These examples are intended to illustrate, not limit, the scope of the claimed invention.

#### Test Extracts and Compounds

In general, novel anti-pathogenic drugs or plant protectants are identified from large libraries of both natural product or synthetic (or semi-synthetic) extracts

or chemical libraries according to methods known in the Those skilled in the field of drug discovery and art. development will understand that the precise source of test extracts or compounds is not critical to the 5 screening procedure(s) of the invention. Accordingly, virtually any number of chemical extracts or compounds can be screened using the methods described herein. Examples of such extracts or compounds include, but are not limited to, plant-, fungal-, prokaryotic- or animal-10 based extracts, fermentation broths, and synthetic compounds, as well as modification of existing compounds. Numerous methods are also available for generating random or directed synthesis (e.g., semi-synthesis or total synthesis) of any number of chemical compounds, 15 including, but not limited to, saccharide-, lipid-, peptide-, and nucleic acid-based compounds. Synthetic compound libraries are commercially available from Brandon Associates (Merrimack, NH) and Aldrich Chemical (Milwaukee, WI). Alternatively, libraries of natural 20 compounds in the form of bacterial, fungal, plant, and animal extracts are commercially available from a number of sources, including Biotics (Sussex, UK), Xenova (Slough, UK), Harbor Branch Oceangraphics Institute (Ft. Pierce, FL), and PharmaMar, U.S.A. (Cambridge, MA). 25 addition, natural and synthetically produced libraries are produced, if desired, according to methods known in the art, e.g., by standard extraction and fractionation methods. Furthermore, if desired, any library or compound is readily modified using standard chemical, 30 physical, or biochemical methods.

In addition, those skilled in the art of drug discovery and development readily understand that methods for dereplication (e.g., taxonomic dereplication, biological dereplication, and chemical dereplication, or any combination thereof) or the elimination of replicates

or repeats of materials already known for their antipathogenic activity should be employed whenever possible.

When a crude extract is found to have antipathogenic activity, further fractionation of the

5 positive lead extract is necessary to isolate chemical constituents responsible for the observed effect. Thus, the goal of the extraction, fractionation, and purification process is the careful characterization and identification of a chemical entity within the crude extract having anti-pathogenic activity. Methods of fractionation and purification of such heterogenous extracts are known in the art. If desired, compounds shown to be useful agents for the treatment of pathogenicity are chemically modified according to methods known in the art.

There now follow examples of high-throughput systems useful for evaluating the efficacy of a molecule or compound in promoting resistance to a pathogen or inhibiting a pathogen. These examples are provided to illustrate, not limit, the invention.

#### Nematode Bioassay System

To enable mass screening of large quantities of natural products, extracts, or compounds in an efficient and systematic fashion, Caenorhabditis elegans L4

25 hermaphrodite larvae are cultured in wells of a microtiter plate, facilitating the semiautomation of manipulations and full automation of data collection. As is discussed above, we have discovered that P. aeruginosa UCBPP-PA14 infects and kills C. elegans, whereas P.

30 aeruginosa UCBPP-PA14 carrying a mutagenized virulence gene is non-pathogenic. If a pathogen has diminished, pathogenicity then L4 worms live, develop into adult hermaphrodites, and produce thousands of live progeny. Accordingly, if C. elegans is incubated with the

pathogen, the worms will die, unless a compound is present to reduce *P. aeruginosa* pathogenicity. The presence of such live progeny is easily detected using a variety of methods, including visual screening with standard microscopes.

To evaluate the ability of a test compound or extract to promote a host's resistance to a pathogen or to repress pathogenicity of a pathogen, a test compound or extract is inoculated at an appropriate dosage into 10 NGM agar seeded with an appropriate amount of an overnight culture of a pathogen, e.g., P. aeruginosa UCBPP-PA14. If desired, various concentrations of the test compound or extract can be inoculated to assess dosage effect on both the host and the pathogen. Control 15 wells are inoculated with non-pathogenic bacteria (negative control) or a pathogen in the absence of a test compound or extract (positive control). Plates are then incubated 24 hours at 37°C to facilitate the growth of the pathogen. Microtiter dishes are subsequently cooled 20 to 25°C, and two C. elegans L4 hermaphrodite larva are added to the plate and incubated at 25°C, the upper limit for normal physiological integrity of C. elegans. At an appropriate time interval, e.g., 4 to 5 days, wells are examined for surviving progeny, e.g., by monitoring 25 motion of worms using a motion detector.

Comparative studies between treated and control larvae are used to determine the relative efficacy of the test molecule or compound in promoting the host's resistance to the pathogen or inhibiting the virulence of the pathogen. A test compound which effectively stimulates, boosts, enhances, increases, or promotes the host's resistance to the pathogen or which inhibits, inactivates, suppresses, represses, or controls pathogenicity of the pathogen and does not adversely affect the normal physiology, reproduction, or

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development of the worms is considered useful in the invention.

#### Plant Bioassay System

To enable mass screening of large quantities of 5 natural products, extracts, or compounds in an efficient and systematic fashion, host plants (e.g., seeds, seedlings, plantlets, embryos, mature plants, or leaves) are cultured in wells of a microtiter plate or any other suitable container, facilitating the semiautomation of 10 manipulations and full automation of data collection. Particular examples of suitable plant hosts useful in this bioassay include, without limitation, petunia, tomato, potato, tobacco, Arabidopsis, soybean, corn, wheat, rye, rice, barley, or any other plant of 15 commercial or agricultural significance. Methods for culturing plants are known in the art (see, e.g., Vasil, I.K., Cell Culture and Somatic Cell Genetics of Plants Vol I, II, III, Laboratory Procedures and Their Applications, Academic Press, New York, 1984; Dixon R.A., 20 Plant Cell Culture - A Practical Approach, IRL Press, Oxford University, 1985). As is discussed above, we have discovered that P. aeruginosa UCBPP-PA14 infects and kills Arabidopsis thaliana, whereas P. aeruginosa UCBPP-PA14 carrying a mutagenized virulence gene is non-25 pathogenic. Accordingly, if a pathogen has diminished pathogenicity, the plant will not develop symptoms or, alternatively, will develop attenuated symptoms relative to control plants. If Arabidopsis thaliana plants are incubated with the pathogen, the plants will die or have 30 a variety of disease symptoms (e.g., chlorosis or softrot), unless a compound is present to reduce P. aeruginosa pathogenicity. The presence of such live seedlings and their associated disease symptoms is easily

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detected using a variety of methods, including visual screening.

To evaluate the ability of a test compound or extract to promote a host's (e.g., Arabidopsis thaliana) 5 resistance to a pathogen or to repress pathogenicity of a pathogen, a test compound or extract is inoculated at an appropriate dosage into a tissue culture media (e.g., a solidified agar-based medium). In addition, if desired, the host plant can be pretreated with the candidate plant 10 protectant or anti-pathogen compound by any conventional means, e.g., a seedling or plantlet can be sprayed with a solution containing the test compound. Host plants are assayed using any standard pathogenesis screening system, e.g., the Arabidopsis and lettuce leaf infiltration 15 assays described above, or by standard vacuum infiltration techniques. For example, host seedlings are vacuum infiltrated with the pathogen according to standard methods. After vacuum infiltration seedlings are cultured according to methods known in the art (e.g., 20 methods for culturing Arabidopsis are found in Methods in Arabidopsis Research, Koncz, C., Chua, N.-H., Schell, J., eds., World Scientific Publishing Co. Pte. Ltd., Singapore, 1992). If desired, various concentrations of the test compound or extract can be inoculated to assess 25 dosage effect on both the host and the pathogen. Control seedlings are infiltrated with non-pathogenic bacteria (negative control) or a pathogen in the absence of a test compound or extract (positive control). At an appropriate time interval, e.g., 3 to 5 days, seedlings 30 are examined for disease symptoms. Comparative studies between treated and control seedlings are used to determine the relative efficacy of the test molecule or compound in promoting the host's resistance to the pathogen or inhibiting the virulence of the pathogen. 35 test compound which effectively stimulates, boosts,

enhances, increases, or promotes the host's resistance to the pathogen or which inhibits, inactivates, suppresses, represses, or controls pathogenicity of the pathogen and does not adversely affect the normal physiology, 5 reproduction, or development of the seedlings is considered useful in the invention.

#### <u>Use</u>

The methods of the invention provide a simple means for identifying microbial virulence factors and 10 compounds capable of either inhibiting pathogenicity or enhancing an organism's resistance capabilities to a Accordingly, a chemical entity discovered to have medicinal or agricultural value using the methods described herein are useful as either drugs, plant 15 protectants, or as information for structural modification of existing anti-pathogenic compounds, e.g., by rational drug design. Such methods are useful for screening compounds having an effect on a variety of pathogens including, but not limited to, bacteria, 20 viruses, fungi, annelids, nematodes, platyhelminthes, and protozoans. Examples of pathogenic bacteria include, without limitation, Aerobacter, Aeromonas, Acinetobacter, Agrobacterium, Bacillus, Bacteroides, Bartonella, Bortella, Brucella, Calymmatobacterium, Campylobacter, 25 Citrobacter, Clostridium, Cornyebacterium, Enterobacter, Escherichia, Francisella, Haemophilus, Hafnia, Helicobacter, Klebsiella, Legionella, Listeria, Morganella, Moraxella, Proteus, Providencia, Pseudomonas, Salmonella, Serratia, Shigella, Staphylococcus, 30 Streptococcus, Treponema, Xanthomonas, Vibrio, and Yersinia.

For therapeutic uses, the compositions or agents identified using the methods disclosed herein may be administered systemically, for example, formulated in a

pharmaceutically-acceptable buffer such as physiological saline. Preferable routes of administration include, for example, subcutaneous, intravenous, interperitoneally, intramuscular, or intradermal injections which provide 5 continuous, sustained levels of the drug in the patient. Treatment of human patients or other animals will be carried out using a therapeutically effective amount of an anti-pathogenic agent in a physiologically-acceptable carrier. Suitable carriers and their formulation are 10 described, for example, in Remington's Pharmaceutical Sciences by E.W. Martin. The amount of the antipathogenic agent to be administered varies depending upon the manner of administration, the age and body weight of the patient, and with the type of disease and 15 extensiveness of the disease. Generally, amounts will be in the range of those used for other agents used in the treatment of other microbial diseases, although in certain instances lower amounts will be needed because of the increased specificity of the compound. A compound is 20 administered at a dosage that inhibits microbial proliferation. For example, for systemic administration a compound is administered typically in the range of 0.1 ng - 10 g/kg body weight.

For agricultural uses, the compositions or agents
identified using the methods disclosed herein may be used
as chemicals applied as sprays or dusts on the foliage of
plants. Typically, such agents are to be administered on
the surface of the plant in advance of the pathogen in
order to prevent infection. Seeds, bulbs, roots, tubers,
and corms are also treated to prevent pathogenic attack
after planting by controlling pathogens carried on them
or existing in the soil at the planting site. Soil to be
planted with vegetables, ornamentals, shrubs, or trees
can also be treated with chemical fumigants for control
of a variety of microbial pathogens. Treatment is

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preferably done several days or weeks before planting. The chemicals can be applied by either a mechanized route, e.g., a tractor or with hand applications. In addition, chemicals identified using the methods of the assay can be used as disinfectants.

All publications and patents mentioned in this specification are herein incorporated by reference to the same extent as if each individual publication or patent was specifically and individually indicated to be incorporated by reference.

From the feregoing description, one skilled in the art can easily ascertain the essential characteristics of this invention, can make various changes and modifications of the invention to adapt it to various usages and conditions. Thus, other embodiments are also within the claims.

#### Deposit

Pseudomonas aeruginosa strain UBCPP-PA14 has been deposited with the American Type Culture Collection on 20 March 22, 1995, and bears the accession number ATCC No. 55664. Applicants acknowledge their responsibility to replace this strain should it loose viability before the end of the term of a patent issued hereon, and their responsibility to notify the American Type Culture 25 Collection of the issuance of such a patent, at which time the deposit will be made available to the public. Prior to that time the deposit will be made available to the Commissioner of Patents under terms of CFR §1.14 and 35 USC §112.

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#### SEQUENCE LISTING

#### (1) GENERAL INFORMATION:

- (i) APPLICANT: Ausubel, Frederick M.
  Rahme, Laurence G.
  Tan, Man-Wah
  Ruvkun, Gary B.
- (ii) TITLE OF INVENTION: METHODS OF SCREENING COMPOUNDS USEFUL FOR PREVENTION OF INFECTION OR PATHOGENICITY
- (iii) NUMBER OF SEQUENCES: 2
- (iv) CORRESPONDENCE ADDRESS:
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  - (E) COUNTRY: USA
  - (F) ZIP: 02110-2804
- (V) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible
  - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30

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- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: US
  - (B) FILING DATE:
  - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Clark, Paul T.
  - (B) REGISTRATION NUMBER: 30,162
  - (C) REFERENCE/DOCKET NUMBER: 00786/263001
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    - (C) TELEX: 200154
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 17 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA (genomic)
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

### GCTAGTAGTC GATGACC

(2) INFORMATION FOR SEQ ID NO:2:

- (2) 200 000 000 000
  - (i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 17 base pairs
    - (B) TYPE: nucleic acid

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- (C) STRANDEDNESS: single
  (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

GCTGGCATCA ACCATGC

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### Claims

- 1. A method for identifying a compound which is capable of inhibiting a pathogen in a eukaryotic organism said method comprising
- (a) exposing at least two different eukaryotic organisms, at least one of said organisms being a non-rodent, to a single pathogen in the presence of at least one candidate compound; and
- (b) identifying a compound that inhibits said 10 pathogen in each of said eukaryotic organisms.
  - 2. The method of claim 1, wherein said eukaryotic organisms include either
    - (a) a vertebrate and a plant;
    - (b) a vertebrate and an invertebrate;
- (c) a plant and an invertebrate;
  - (d) two vertebrates of different genera; or
  - (e) two invertebrates of different genera.
  - 3. The method of claim 1, wherein said eukaryotic organisms include two plants of different genera.
- 20 4. The method of claim 2, wherein said vertebrate is a mammal.
  - 5. A method for identifying a compound which is capable of inhibiting a pathogen in a non-rodent eukaryotic host organism, comprising
- 25 (a) exposing said a non-rodent eukaryotic host organism to a single pathogen in the presence of at least one candidate compound; and
  - (b) identifying a compound that inhibits the pathogen in said eukaryotic host organism.

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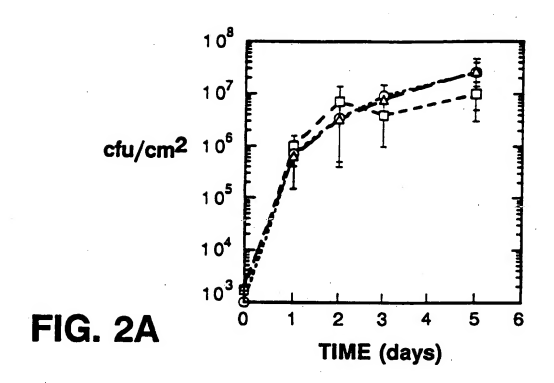
- 6. The method of claims 1 or 5, wherein said pathogen is a bacterium.
- 7. The method of claim 6, wherein said bacterium is Pseudomonas aeruginosa UCBPP-PA14.
- 5 8. The method of claims 1 or 5, wherein said non-rodent eukaryotic host organism is a nematode.
  - 9. The method of claim 8, wherein said nematode is Caenorhabditis elegans.
- 10. The method of claims 1 or 5, wherein said 10 said non-rodent eukaryotic organism is *Arabidopsis*.
  - 11. A method for identifying a pathogenic virulence factor, comprising
- (a) identifying a pathogen which is capable of infecting at least two different eukaryotic organisms, at
   15 least one of said organisms being a non-rodent;
  - (b) generating a mutant of said pathogen;
  - (c) exposing each of said organisms to said
    mutated pathogen;
- (d) determining whether said mutated pathogen is 20 capable of causing disease in each of said organisms, a reduction of disease in both of said organisms relative to that caused by said wild-type pathogen indicating a mutation in said pathogenic virulence factor; and
- (e) using said mutation as a marker for 25 identifying said pathogenic virulence factor.
  - 12. A method for mutating a pathogenic virulence factor, comprising

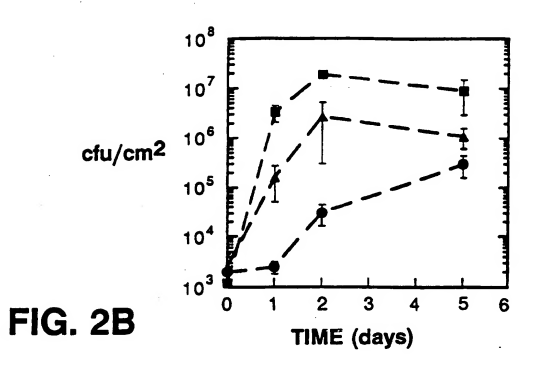
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- (a) identifying a pathogen which is capable of infecting at least two different eukaryotic organisms, at least one of said organisms being a non-rodent;
  - (b) generating a mutant of said pathogen;
- (c) exposing each of said organisms to said mutated pathogen; and
- (d) determining whether said mutated pathogen is capable of causing disease in each of said organisms, a reduction of disease in both of said organisms relative
   to that caused by said wild-type pathogen indicating a mutation in said pathogenic virulence factor.
  - 13. A method of reducing the virulence of a pathogen, comprising
- (a) identifying a pathogen which is capable of 15 infecting at least two different eukaryotic organisms, at least one of said organisms being a non-rodent;
  - (b) generating a mutant of said pathogen;
  - (c) exposing each of said organisms to said mutated pathogen; and
- 20 (d) determining whether said mutated pathogen is capable of causing disease in each of said organisms, a reduction of disease in both of said organisms relative to that caused by said wild-type pathogen indicating a reduction in said pathogen virulence.

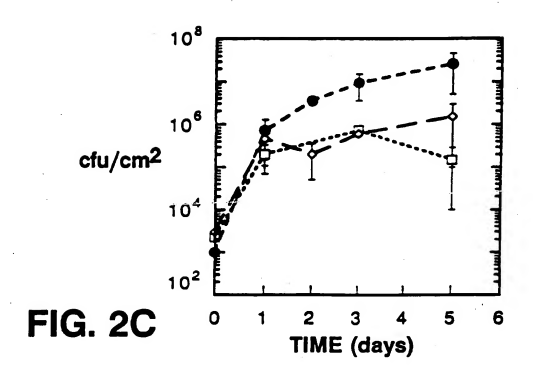


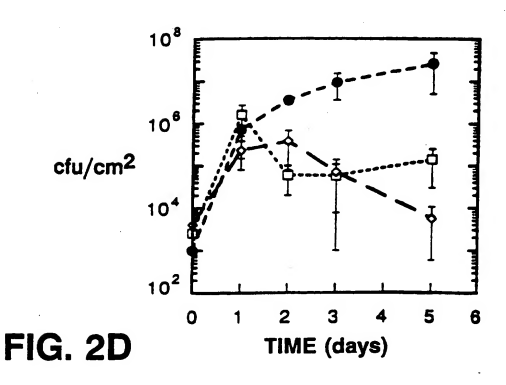
FIG. 1





# SUBSTITUTE SHEET (RULE 26)





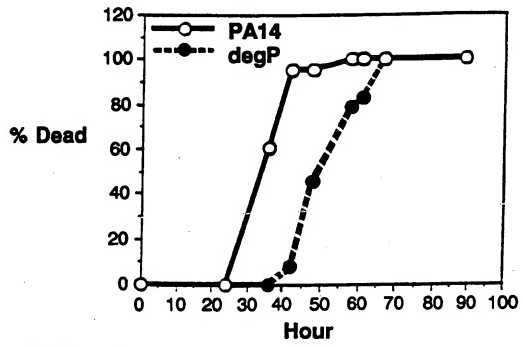


FIG. 3

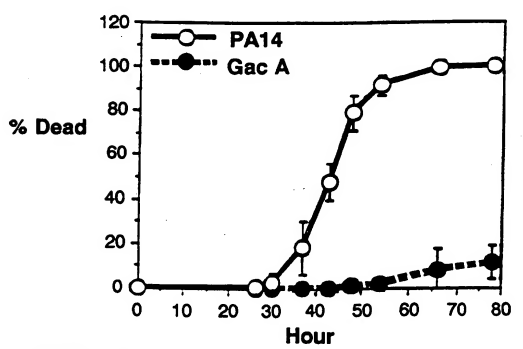


FIG. 4

# SUBSTITUTE SHEET (NULE 26)

## INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04210

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) :A61K 49/00; C12Q 1/18; C12N 15/00  US CL :435/32, 172.3; 424/9.2				
According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIELDS SEARCHED				
Minimum do	ocumentation searched (classification system followers	ed by classification symbols)		
U.S. : 435/32, 172.3, 253.3; 424/9.2; 935/76				
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)				
APS, CAS Search terms: in vivo, screen?, test?, pathogen?, infect?, bacter?, host, virulence, mutat?, arabidopsis, plant, nematode				
C. DOCU	JMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.	
	US, A, 4,713,378 (PERRONE ET column 11, lines 21-68, column 1		1, 2, 4-6	
	US, A, 5,366,995 (SAVAGE ET column 3, lines 18-30, column 8,		1, 3, 5, 6, 10	
	Journal of Bacteriology, Volume 1 1988, Xu et al, "Molecular Cloni Virulence in <i>Pseudomonas solana</i> see entire document.	ng of Genes That Specify	11-13	
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Further	r documents are listed in the continuation of Box C	See patent family annex.		
Special categories of cited documents:  'A' document defining the general state of the art which is not considered		"T" later document published after the inte- date and not in conflict with the applica principle or theory underlying the inve	tion but cited to understand the	
to be of particular relevance  *E* earlier document published on or after the international filing date		"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step		
*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other		when the document is taken alone  "Y" document of particular relevance; the claimed invention cannot be		
special reason (as specified)  "O" document referring to an oral disclosure, use, exhibition or other means		considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art		
*P* document published prior to the international filing date but later than the priority date claimed		*&* document member of the same patent family		
Date of the ac	stual completion of the international search	Date of mailing of the international sea	rch report	
24 JUNE 1996		0 9 JUL 1996		
Name and mailing address of the ISA/US  Commissioner of Patents and Trademarks		Authorized officer		
Box PCT Washington, D.C. 20231		PRASAD MURTHY AUGUSTIC		
Facsimile No. (703) 305-3230		Telephone No. (703) 308-0196		

# INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04210

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)			
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:			
1.		Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
2.		Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:	
3.		Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box	× II (	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
This International Searching Authority found multiple inventions in this international application, as follows:			
Please See Extra Sheet.			
1.	X	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.	
2.		As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	
3.		As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:	
4.		No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	
R	emark	on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.	

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04210

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

Group I, claim(s) 1-10, drawn to screening assays for compounds which inhibit pathogens.

Group II, claim(s) 11-13, drawn to method of identifying and mutating virulence genes.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: the special technical feature of the Group I invention is the identification of compounds which inhibit pathogens in vivo, while the special technical feature of the Group II invention is the mutation and subsequent identification of virulence genes of a pathogen. Since the special technical feature of the Group I invention is not present in the Group II claims and the special technical feature of the Group II invention is not present in the Group I claims, unity of invention is lacking.